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FILE COVERS 1907 - 30 Nov 2008 VOL 149 ISS 23 FILE LAST UPDATED: 28 Nov 2008 (20081128/ED)

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=> s 11 and hexadiene 8039 HEXADIENE L2 8 L1 AND HEXADIENE

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=> d 1-8 bib abs
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- ANSWER 1 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN
- 2006:1187835 CAPLUS AN
- DN 146:122117
- TΙ Synthesis and Reactivity of Azapalladacyclobutanes
- Ney, Joshua E.; Wolfe, John P. AU
- CS Department of Chemistry, University of Michigan, Ann Arbor, MI, 48109-1055, USA
- SO Journal of the American Chemical Society (2006), 128(48), 15415-15422 CODEN: JACSAT: ISSN: 0002-7863
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 146:122117
- GT

AB N-Sulfonvl aziridines undergo oxidative addition to palladium(0) complexes yielding azapalladacyclobutane amido-complexes, which rearrange to azapalladabicyclo[3.2.1]octanes via copper-catalyzed carbopalladation. Palladium(0) species generated in situ from Pd2(dba)3 and 1,10-phenanthroline undergo oxidative addition of 2-methyl-1-nosyl- and 2-(4-R-3-butenyl)-1-tosylaziridines (R = H, BuO2C), affording 3-Me- or 3-(RCH:CHCH2CH2)-1-X-2-(phen)-1,2azapalladacyclobutanes (15, 8, 16, resp., same R; X = 4-NO2C6H4SO2, 4-MeC6H4SO2), whereas aziridines containing less electroneg. or saturated alkyl groups show no reaction. The resulting azapalladacyclobutane complexes undergo intramol. carbopalladation in the presence of copper(I) iodide to afford azapalladabicyclo[3.2.1]octanes (I, 17, R = H; 18, R = CO2Bu). A deuterium-labeling experiment indicates that the oxidative addition proceeds

via

SN2-type attack of palladium(0) on the less-hindered carbon of the aziridine ring and that alkene insertion occurs in a syn-fashion. The azapalladabicyclo[3.2.1]octane complexes undergo oxidative palladium-carbon bond functionalization in the presence of copper(II) bromide.

RE.CNT 136 THERE ARE 136 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- ANSWER 2 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN L2
- AN 2004:390260 CAPLUS
- DN 140:391378
- TΙ Method for preparation of compounds containing palladium (0) and alkadienes
- IN Kayser, Bernd; Karch, Ralf; Briel, Oliver; Kleinwaechter, Ingo
- PA Umicore Ag & Co. Kg, Germany
- SO PCT Int. Appl., 27 pp.
- CODEN: PIXXD2
- Patent T.A German
- FAN.CNT 1

PATENT NO.

KIND DATE

APPLICATION NO.

DATE

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WO 2004039819 A2 20040513 WO 2003-EP12085
WO 2004039819 A3 20041021
PΤ
                                                                        20031030
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
              CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
              GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
              LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO,
              NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ,
              TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
          RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
              BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE,
              ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK,
              TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     DE 10250901
                           A1
                                 20040519 DE 2002-10250901 20021031
     CA 2504418
                           A1
                                  20040513 CA 2003-2504418
                                                                        20031030
     AU 2003287975
                           A1
                                  20040525 AU 2003-287975
                                                                        20031030
     EP 1558621
                           A2 20050803
                                              EP 2003-779823
                                                                        20031030
     EP 1558621
                                 20080123
                           B1
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
              IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
CN 1720253 A 20060111 CN 2003-80104632 

JP 2006504765 T 20060209 JP 2004-547623 

AT 384730 T 20080215 AT 2003-779823 

RU 2334754 C2 20080257 RU 2005-116671 

2A 2005003426 A 20060927 ZA 2005-3426 

IN 2005KN00755 A 20060099 IN 2005-EN755 

US 20060106239 A1 2006018 US 2005-533492 

PRAI DE 2002-10250901 A 20021031 

WO 2003-EP12085 W 20031030
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                                                                        20031030
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                                                                         20050428
                                                                        20051003
     CASREACT 140:391378; MARPAT 140:391378
OS
AB
     The invention relates to a method for the production of a compound containing
     palladium(0), which consists in reacting a palladium
     compound with one or several compds. of general R1R2C:C(R3) {(A)x}C(R6):CR4R5
     (R1-R6 = halo, alkoxy, CN, NO2, amino, organocarbonyl, carboxy, amido,
     substituted alkenyl, organosilyl, organosulfonyl, organothio,
     organophosphino, etc.; A = organoalkenyl, O, S, organoamino, organosilyl,
     substituted ring, x = 2-4), cyclic [Si(R12)(C13CR14R15)0]n (n = 3-20, R12
     = H, OH, (un)substituted C1-20 alkyl, OC1-10 alkyl, etc.; R13-R15 = halo,
     CN, NO2, alkyl, organoamino, organocarbonyl, organocarboxy, organoamido,
     substituted alkenyl, organosilyl, organosiloxy, organothio,
     organosulfonvl, organophosphino, etc.), or
     Term-O-\{Si(R16)(CR19CR17R18)O\}v\{Si(R20)2O\}w\}-Term (v, w = 1-1000; v+w =
     0-1000; R16, R20 = H, OH, (un) substituted C1-10 alkyl, etc.; R17-R19 =
     halo, CN, NO2, alkyl, organoamino, organocarbonyl, organocarboxy,
     organoamido, substituted alkenyl, organosilyl, organosiloxy, organothio,
     organosulfonyl, organophosphino, etc.) in the presence of a base. Thus,
     reaction of sodium tetrachloropalladate with diallyl ether in presence of
     sodium hydrogen carbonate in MeOH gave 81% title compound in 4h. The
     compds. containing palladium(0) are suitable for use as
     homogeneous catalysts, as precursors in the in situ production of homogeneous
     catalysts or as precursors in the production of heterogeneous catalysts.
     ANSWER 3 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN
AN
     1996:251769 CAPLUS
     125:10316
OREF 125:2273a,2276a
TI
     Palladium(0)-catalyzed phenoxycarbonylation of allylic
     carbonates
ΆΠ
     Goux, Catherine; Lhoste, Paul; Sinou, Denis; Masdeu, Anna
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Laboratoire de Synthese Asymetrique, URA 463 du CNRS, CPE Lyon, Universite

Claude Bernard Lyon I, Villeurbanne, 69622, Fr.

Journal of Organometallic Chemistry (1996), 511(1-2), 139-43

CS

SO

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CODEN: JORCAI; ISSN: 0022-328X
PB
    Elsevier
DT
    Journal
LA English
AB Phenoxycarbonylation of various allylic carbonates under various
    conditions in THF is described. The nature and ratio of the products
     formed are dependent on the presence of water, carbon monoxide pressure
     and addition of various inorg, halides. The formation of a product arising
     from dimerization of the allylic carbonate is discussed. The
     phenoxycarbonylation of (E)-cinnamyl Et carbonate gave
     (E)-4-phenyl-3-butenoic acid Me ester and cinnamyl Ph ether.
   ANSWER 4 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN
AN
    1993:212686 CAPLUS
DN
   118:212686
OREF 118:36647a,36650a
TI Palladium-catalyzed coupling of aryl halides, non-conjugated dienes and
    carbon nucleophiles
IN
     Larock, Richard C.
PA
     Iowa State University Research Foundation Inc., USA
SO
    U.S., 6 pp.
    CODEN: USXXAM
DT
    Patent
LA
    English
FAN.CNT 1
                                         APPLICATION NO.
                       KIND DATE
     PATENT NO.
                                                                 DATE
    US 5164518
                               19921117
                                         US 1991-729428
PΙ
                        A
                                                                  19910712
    WO 9301172
                        A1
                               19930121
                                          WO 1992-US5753
                                                                  19920709
        W: JP
        RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, MC, NL, SE
PRAI US 1991-729428
                     A 19910712
    CASREACT 118:212686; MARPAT 118:212686
OS
AB RCH:CR1(CHR1)pCR1:CHR2 (R - R2 = H, alkyl, aryl, aralkyl; RR2 = alkylene;
     p = ≥1) were subjected to the title reaction to give
     RR3CH(CHR1)pCR1:CR1CHR2CR4R5R6 (R3 = aryl; R4 = H, alkyl, aryl, aralkyl;
     R5, R6 = cyano, acyl, alkoxycarbonyl, arylsulfonyl; R5R6 = oxoalkylene,
     carbonyloxyalkylene). Thus, CH2:CHCH2CH2CH2CH2 was treated with PhI and
     CH2(CO2Et)2 in the presence of bis(dibenzylideneacetone)palladium
     (0), Bu4NCl, and NaHCO3 to give 82%
     (E)-Ph(CH2)3CH:CHCH2CH(CO2Et)2.
L2 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN
AN
    1992:129191 CAPLUS
DN
    116:129191
OREF 116:21887a,21890a
ΤI
    (R2PC2H4PR2)Pd0 alkene and ethyne complexes
    Krause, Jochen; Bonrath, Werner; Poerschke, Klaus R.
AU
CS
    Max-Planck-Inst. Kohlenforsch., Muelheim ander Ruhr, D-4330, Germany
    Organometallics (1992), 11(3), 1158-67
SO
     CODEN: ORGND7; ISSN: 0276-7333
DT
    Journal
LA
    English
OS
    CASREACT 116:129191
    The palladium n3-allyl complex Pd(n3-C3H5)2 reacts with bidentate
     phosphines iso-Pr2PC2H4PPr-iso2 and tert-Bu2PC2H4PBu-tert2) below
     -30° to yield the light yellow, microcryst. palladium(II)
     \eta1-allyl compds. (R2PC2H4PR2)Pd(\eta1-C3H5)2 (R = iso-Pr (I), tert-Bu
     (II), which are stable to about -30°. Above 30°, the allyl
     substituents of I and II couple with reduction of palladium to form a mixture
of
    (R2PC2H4PR2)Pd0 1,5-hexadiene complexes. When this reaction is
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carried out in 1,5-hexadiene, the complex $(R2PC2H4PR2)Pd(\eta2-C6H10)$ R = iso-Pr (III), tert-Bu (IV) are obtained in pure form. According to IR and NMR spectral data, the palladium atoms in III and IV are coordinated at low temperature by the chelating diphosphane and (statically) by one of the 2 diene double bonds in a trigonal-planar geometry. At higher temps., a rapid exchange of the coordinated and uncoordinated double bonds occurs, passing through an intermediated with C2 symmetry. When suspension of I in pentane and II and THF are warmed to 20°, dinuclear diastereomers rac-meso-{(R2PC2H4PR2)Pd}2 (μ-η2:η2-C6H10) are obtained, which upon treatment with 1.5hexadiene furnish mononuclear derivs. III and IV. Similarly, when I is reacted with 1,5-cyclooctadiene at 20°, the mono- and dinuclear interconvertible complexes (iso-Pr2PC2H4PPr-iso2)Pd(n2-C8H12) and $\{(iso-Pr2PC2H4PPr-iso2)Pd\}2(\mu-\eta2:\eta2-C8H11) \text{ are produced. From }$ the reaction of I and II with ethene, stable complexes (R2PC2H4PR2)Pd(C2H4) (V), result. The colorless mononuclear palladium(0) ethyne complexes (R2PC2H4PR2)Pd(C2H2) may be prepared (a) by a displacement reaction of one of the isolated alkene complexes with ethyne, (b) by a reductive elimination and concomitant displacement reaction of the n1-allyl complexes I and II with ethyne, or (c) in a one-pot synthesis from either Pd(m3-C3H5)2 or Pd(n3-2-MeC3H4)2 with iso-Pr2PC2H4PPr-iso2 or tert-Bu2PC2H4PBu-tert2, resp., and ethyne in excess. When the mononuclear ethyne complexes V are combined with an equimolar amount of the corresponding \(\eta 1 - allyl \) complex I or II, or the alkene complexes, yellow dinuclear palladium(0) complexes {(R2PC2H4PR2)Pd]2(µ-C2H2) are produced, in which the ethyne ligand bridges 2 Pd atoms. All the diphosphine) palladium(0) alkene and ethyne complexes exhibit a trigonal-planar coordination geometry about the Pd atom. Most of the compds. have been isolated in high yield.

- L2 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 1988:630210 CAPLUS
- DN 109:230210

OREF 109:38064h,38065a

- IT Fluoroalkylation and fluoroalkoxylation. Part 28. Palladium(
 0)-induced addition of fluoroalkyl iodides to alkenes: an electron-transfer process.
- AU Chen, Qingyun; Yang, Zhenyu; Zhao, Chengxue; Qiu, Zaiming
- CS Shanghai Inst. Org. Chem., Acad. Sin., Shanghai, Peop. Rep. China
- SO Journal of the Chemical Society, Perkin Transactions 1: Organic and Bio-Organic Chemistry (1972-1999) (1988), (3), 563-7 CODEN: JCPRB4; ISSN: 0300-922X
- DT Journal LA English

AB

OS CASREACT 109:230210

involved. Also, the formation of

readily with alkenes under mild conditions to give the corresponding adducts in high yields. Dichlorobis(triphenylphosphine)palladium or fluoroalkylbis(triphenylphosphine)palladium iodide fails to induce the reaction under the same conditions.

Diiodobis(triphenylphosphine)palladium, but not the fluoroalkylated complex of Pd, is formed in the reaction confirmed by XPS. The reaction can be partially suppressed by radical inhibitors, such as 2-nitro-2-nitrosopropane or di-tert-Bu nitroxide, but no effects are observed by the presence of oxygen in air of p-dinitrobenzene [molar ratio p-DNB:PdO = 1:1] unless an excess of p-DNB(10:1) was added. In the presence of diallyl ether, fluoroalkylated THF derivs. are formed. All these results indicate that a radical-chain process initiated by single electron transfer from palladium(0) to iodide might be

In the presence of catalytic amts, of (Ph3P)4Pd, fluoroalkyl iodides react

fluoroalkylbis(triphenylphosphine)palladium iodide is shown to proceed through a radical intermediate by ESR trapping techniques.

- ANSWER 7 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN 1.2
- AN 1979:438309 CAPLUS
- DM 91:38309
- OREF 91:6256h,6257a
- (Polystyrylbipyridine)palladium(0): heterogeneous
 - catalyst for the mild hydrogenation of dienes to monoenes
- ΑU Card, Roger J.; Neckers, Douglas C.
- CS Dep. Chem., Bowling Green State Univ., Bowling Green, OH, 43403, USA
- SO Israel Journal of Chemistry (1979), Volume Date 1978, 17(4), 269-73
 - CODEN: ISJCAT; ISSN: 0021-2148
- DT Journal
- LA English
- AB Partial hydrogenation of dienes using the title catalyst could be effected in swelling solvents, such as THF or benzene under 1 atm H2 and 25°. The hydrogenation of hexadiene isomers results in the formation of 1-, 2- and 3-hexene, with <1% hexane formation up to consumption of the 1st equivalent H2. Cyclohexadiene undergoes H transfer to form cyclohexene and benzene.
- ANSWER 8 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 1975:111508 CAPLUS
- DN 82:111508
- OREF 82:17813a,17814a
- Palladium(0)-catalyzed mercury extrusion.
 - Stereoselective conversion of bis(propenyl)mercury into 2,4hexadiene
- AU Vedejs, E.; Weeks, P. D.
- CS Dep. Chem., Univ. Wisconsin, Madison, WI, USA
- SO Tetrahedron Letters (1974), (36), 3207-10
- CODEN: TELEAY; ISSN: 0040-4039
- Journal DT LA
- English
- AR (Ph3P) 4Pd induced stereoselective metallic Hg extrusion from bis(propenyl)mercury to form 2,4-hexadienes. Other Pd(II) complexes induced diene formation but Hg was not extruded as the metal. (PhCH:CH2)2PdCl2 and (MeCH:CH)2Hq, underwent a Heck reaction giving PhCH:CHCH:CH2 and 1-benzyl-3-methyl-π-allylpalladium chloride dimer.

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